(4) Systems that ensure access to information that facilitates timely followup and inspection by FDA.

§803.18 Files.

- (a) User facilities and manufacturers shall establish and maintain MDR event files. All MDR event files shall be prominently identified as such and filed to facilitate timely access.
- (b)(1) For purposes of this part, "MDR event files" are written or electronic files maintained by user facilities and manufacturers. MDR event files may incorporate references to other information, e.g., medical records, patient files, engineering reports, etc., in lieu of copying and maintaining duplicates in this file. MDR event files must contain:
- (i) Information in the possession of the reporting entity or references to information related to the adverse event, including all documentation of the entity's deliberations and decisionmaking processes used to determine if a device-related death, serious injury, or malfunction was or was not reportable under this part.
- (ii) Copies of all MDR forms, as required by this part, and other information related to the event that was submitted to FDA and other entities (e.g., a distributor or manufacturer).
- (2) User facilities and manufacturers shall permit any authorized FDA employee during all reasonable times to access, to copy, and to verify the records required by this part.
- (c) User facilities shall retain an MDR event file relating to an adverse event for a period of 2 years from the date of the event. Manufacturers shall retain an MDR event file relating to an adverse event for a period of 2 years from the date of the event or a period of time equivalent to the expected life of the device, whichever is greater. MDR event files must be maintained for the time periods described in this paragraph even if the device is no longer distributed.
 - (d) [Reserved]
- (e) The manufacturer may maintain MDR event files as part of its complaint file, under §820.198 of this chapter, provided that such records are prominently identified as MDR reportable events. A report submitted under

this subpart A shall not be considered to comply with this part unless the event has been evaluated in accordance with the requirements of §§ 820.162 and 820.198 of this chapter. MDR files shall contain an explanation of why any information required by this part was not submitted or could not be obtained. The results of the evaluation of each event are to be documented and maintained in the manufacturer's MDR event file.

§ 803.19 Exemptions, variances, and alternative reporting requirements.

- (a) The following persons are exempt from the reporting requirements under this part.
- (1) An individual who is a licensed practitioner who prescribes or administers devices intended for use in humans and who manufactures or imports devices solely for use in diagnosing and treating persons with whom the practitioner has a "physician-patient" relationship.
- (2) An individual who manufactures devices intended for use in humans solely for such person's use in research or teaching and not for sale, including any person who is subject to alternative reporting requirements under the investigational device exemption regulations, parts 812 and 813 of this chapter, which require reporting of all adverse device effects.
- (3) Dental laboratories, or optical laboratories.
- (b) Manufacturers or user facilities may request exemptions or variances from any or all of the reporting requirements in this part. The request shall be in writing and include information necessary to identify the firm and device, a complete statement of the request for exemption, variance, or alternative reporting, and an explanation why the request is justified.
- (c) FDA may grant in writing, to a manufacturer or user facility, an exemption, variance or alternative from, or to, any or all of the reporting requirements in this part and may change the frequency of reporting to quarterly, semiannually, annually, or other appropriate time period. These modifications may be initiated by a request as specified in this section, or at the discretion of FDA. When granting